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Assessing the performance of the Cretan Sea ecosystem model with the use of high frequency M3A buoy data set

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Abstract. During the Mediterranean Forecasting System Pilot Project a buoy was deployed in the Cretan Sea and for the first time high-frequency physical and biogeochemical data were collected over an extended period, providing a unique opportunity for the evaluation of an ecosystem model. The model both tuned and validated in the Cretan Sea in the past, is explored and quantified. In addition, the optimal parameter set is determined while the effects of high-frequency forcing are explored. The model results are satisfactory, especially at the upper part of the water column, while the inability of 1-D modelling in fully exploring the hydrodynamics of the particular area is depicted and further developments are suggested.

Key words. Oceanography; general (numerical modeling) – Oceanography; biological and chemical (ecosystems and ecology)

1 Introduction

Heavily populated coastal areas bound the Mediterranean Sea. The economies of these regions are highly dependent on fishing, transportation, recreation and other industries, which in turn depend on a healthy coastal environment. Predicting the behaviour of the marine environment is an essential part of the management of marine resources under anthropogenic stress. Therefore, It is an essential requirement for the marine science community to determine the potential time-scales of predictability of the marine ecosystem if an operational coastal ocean environmental monitoring and forecast system is to be developed. Such a system would provide estimates of the changes in both the physical and the biogeochemical marine environments and an enhanced understanding of the marine ecosystem, essential to guiding resource management. Additionally, it would form an early warning system of potentially harmful ecological events and aid the formulation of cost-effective preventive and remedial

measures. Towards these issues an existing ecosystem model both tuned and validated in the Cretan Sea (Triantafyllou et al., 2002), is explored and its performance is quantified. As a first step historical and recent data sets are used to exploit the model to determine the optimal parameter set. The particular ecosystem model was chosen due to its generic nature in response to the physicochemical environment within which it is placed. With all significant biological pathways in the modelled system included, the model can respond to the physical and chemical forcing in a way that is at least qualitatively correct under a wide range of conditions.

In this study, high-frequency forcing data from the POSEIDON system (Soukissian et al., 1999) and the M3A buoy are used and compared with climatological forcing, in an attempt to investigate whether the performance of the ecosystem model is enhanced. High-frequency forcing model results are further validated with in situ data of nutrients and chlorophyll collected during maintenance trips. Finally, high-frequency data of chlorophyll, nitrate, oxygen and temperature during M3A deployment is hindcast with model results.

2 Materials and methods

2.1 Study area

The area under study is the M3A mooring site located at 35°07'07.54" N, 24°59'44.66" E in the Cretan Sea, north of Heraklion, with a depth of 1100 m (Fig. 1). The open character of the area in conjunction with the significant distance from the north coast of Crete (21 miles) ensures that there is no influence from land activities in the functioning of the specific ecosystem. The simulation period is the M3A deployment period starting from 30 January 2000 until 20 April 2001.

The Cretan Sea at the eastern part of the Mediterranean is dominated by multiple scale circulation patterns. The hydrological structure is complex and is characterised by mesoscale variability. It is an area of deep-water formation

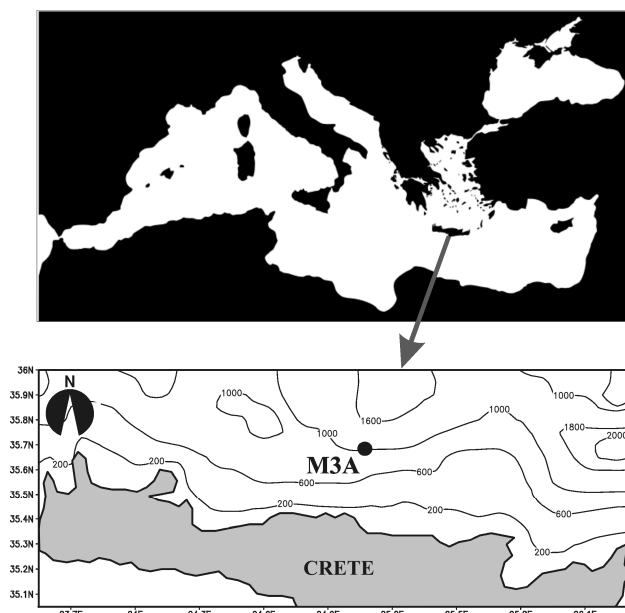


Fig. 1. Map of M3A mooring.

with a sporadic and episodic formation of intermediate water occurring predominantly in the late winter. From spring through to late autumn/ early winter the region is thermally stratified to a depth of 50–70 m. The ecosystem of the outer Cretan Sea has two different modes of operation. During periods of stratification it exhibits a microbial food web, when small phytoplankters dominate, taking advantage of their superior surface-to-volume ratio and a considerable portion of the total organic pool is in the form of detritus. Phosphorus is thought to be the limiting nutrient for phytoplankton and bacterial growth (Becacos-Kontos, 1977; Berland et al., 1980; Krom et al., 1991, 1992; Thingstad and Rassoulzadegan, 1995) with very low concentrations even below the euphotic zone at 200 m. Nutrients are recycled in the top layers with a prominent *P* limitation of both phytoplankton and bacteria, and high surface concentrations and vertically decreasing gradients of DOP and DOC. During this mode very little energy is passed to the higher trophic layers and to the benthos. When the thermocline erodes the system switches into the classical mode during which trapped nutrients find their way to the euphotic zone, initiating a new production bloom and biomass distribution, being linearly ordered according to trophic levels along which the organisms increase sequentially in size (Smetacek and Pollehne, 1986).

The variability observed in biogeochemical parameters is due to the complex hydrological structure characterised by eddies interconnected with jets without defined time scales (Balopoulos, 1996). These features are important because they can transport, entrap and disperse chemicals, particulate matter, small organisms, heat, etc., while they significantly modify the vertical mixing patterns in an area, creating among other phenomena zones of upwelling and downwelling that in turn can have a major effect on the biological

productivity (Krom et al., 1993). Those structures, in conjunction with the presence of transient Mediterranean water (TMW), a mass rich in nutrients and poor in oxygen, can drastically alter the chemical conditions of the intermediate layers of the entire south Aegean (Theocharis et al., 1999), acting as a nutrient source to the euphotic zone or as sinks for organic material to the deeper layers.

2.2 M3A data set

One of the innovations of the MFSP project and the deployment of the M3A buoy was the collection for the first time of high-frequency physical and biogeochemical data over an extended period in the outer Cretan Sea. Previous studies in this region were conducted on a seasonal time-scale, which, in the case of oligotrophic systems, may prove to be inadequate in revealing the ecosystem dynamics since the temporal scales must be considered according to the biological time scales of growth (or generation time), which are rather short (Abbott, 1993). The M3A buoy provides 3-h measurements of temperature, chlorophyll-*a*, nitrate and oxygen, at various depths giving a unique opportunity for hindcasting modelling of the Cretan Sea ecosystem. The locations of the CTD and CT sensors were at 40, 60, and 90 m at line 2 and at 150, 250, 350 and 500 m at line 1. The chlorophyll-*a* and oxygen sensors were deployed at 40, 65, 90 and 115 m and the nitrate sensor at 45 m. A detailed description of the structure and setup of the three moorings comprising the M3A array are presented in Nittis et al. (2003). Chlorophyll-*a*, dissolved oxygen and nitrate data were post-calibrated using *in situ* bottle measurements. The calibration coefficients for the M3A sensors were calculated separately for each period between maintenance cruises, using the reference data collected during each re-deployment when all sensors had been cleaned (Nittis et al., 2003).

2.3 Ecosystem model

In this study the model used is a highly portable coupled physical biogeochemical water column model (Allen et al., 1998) adapted for the oligotrophic Cretan Sea (Triantafyllou et al., 2002). The biogeochemistry is based on the European Regional Seas Ecosystem Model (ERSEM) (Baretta et al., 1995), while the physics are provided by a one-dimensional version of the Princeton Ocean Model (Blumberg and Mellor, 1978).

A turbulence closure model (Mellor and Yamada, 1982) determines the vertical temperature, turbulent kinetic energy, and diffusion coefficient profiles generated by a surface heat flux, salinity and wind stress. Heat transfer is assumed to be via vertical diffusion processes. A background viscosity parameterises other mixing processes (e.g. internal waves). The physical sub-model, in addition, to transport describes irradiance, temperature, and sea surface boundary conditions. The solar radiation is provided to the biogeochemical model as photosynthetically available radiation (PAR) penetrating the water column and becoming extinct with increas-

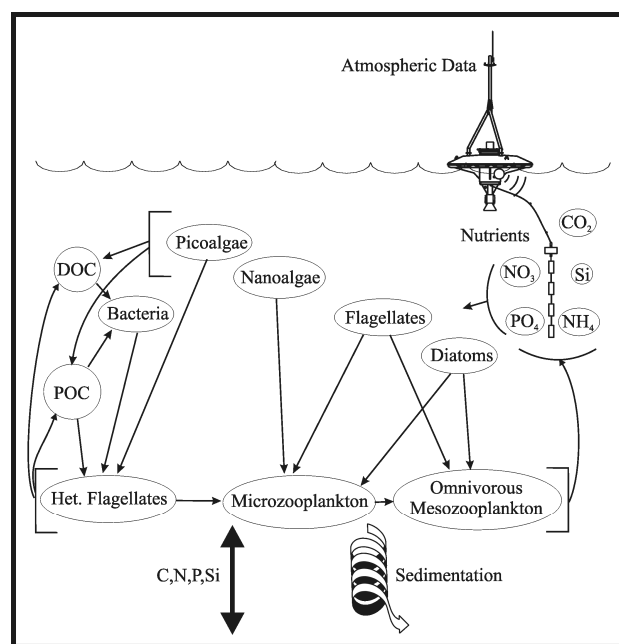


Fig. 2. Ecosystem model food web.

ing depth, according to an extinction coefficient calculated at each time step from the concentrations of primary producers, detritus, bacteria and inorganic suspended matter (Zavatarelli et al., 2000).

The basis of the biogeochemical model consists of state variables with a functional group, approach where organisms are grouped according to their trophic level subdivided according to size classes or feeding method. Two sub-models describe the ecosystem, one for the pelagic and another one for the benthic ecosystem, coupled in terms of carbon and nutrients. The pelagic sub-model includes all the important components, such as nutrient pools, oxygen, phytoplankton, mesozooplankton, microzooplankton, bacteria and detritus. Although the full benthic sub-model depicts a rather diverse system, in this study, due to the depth (> 1000 m) and the oligotrophic nature of the system, benthic processes are described by the simple first order benthic returns module. According to this the detritus falling into the benthos is added in both the particulate and dissolved benthic pools from which a proportion of the total carbon (5% of the particulate and 10% of the dissolved fraction) is remineralised and returned to the water column as CO_2 . The nutrient components of these forms of detritus follow the same proportions with 10% of the remineralised nitrogen added to the nitrate and the rest 90% to the ammonium pools. The remineralised phosphorus is fluxed to the inorganic phosphate pool.

The model food web is modified from the standard ERSEM version 11 (Baretta et al., 1995), according to the characteristics of the ecosystem under study as they have been revealed from literature and analysis of in situ data (Azov, 1991; Stergiou et al., 1997; Tselepidis and Polychronaki, 1996) (Fig. 2). Thus, phytoplankton is com-

posed of four groups with diatoms and flagellates being grazed by microzooplankton and omnivorous mesozooplankton, nanoalgae grazed by microzooplankton, while picoalgae and bacteria are grazed by heterotrophic nanoflagellates. To take into account the importance of bacterial nutrient limitation and the role of dissolved organic material in this oligotrophic system, a modified version (Allen et al., 2002) of the standard ERSEM pelagic sub-model bacteria (Baretta-Bekker et al., 1998; Baretta-Bekker et al., 1995) was used.

The 1-D Cretan Sea ecosystem model (Triantafyllou et al., 2002) which was used has a vertical resolution of 40 layers and total depth of 1100 m, with a finer resolution at the euphotic zone (0–150 m every 10 m). It was forced with six hourly wind speed, humidity, cloud cover and air temperature data obtained from ECMWF. Incident sea surface radiation is calculated from the latitude modified by the cloud cover data using the methods of Patsch (1994), while the extinction coefficient was parameterised according to the measured depth of 1% incident solar irradiance (80–90 m) (Gotsis-Skretas et al., 1999; Ignatiades et al., 1995; Psarra et al., 2000). In this study the model was also forced with high-frequency wind speed and air temperature data for the computation of wind stresses (Hellermann and Rosenstein, 1983) and relaxed to mean monthly sea surface temperature and salinity obtained from a POSEIDON buoy (Soukissian et al., 1999) located 10 miles south of M3A during the period of the M3A deployment (January 2000–April 2001).

2.4 Optimal parameter set

The optimal parameter set for the autotrophic and heterotrophic groups in the model are given in Tables 1 and 2, respectively. The maximum specific uptake rate in all phytoplankton groups was lowered to avoid excess production of dissolved organic carbon (DOC) and subsequent growth of bacteria, which will compete for nutrients with phytoplankton. Faster rates of small cells are, however, retained. Oligotrophic phytoplankton was shown to grow at rates of $0.5\text{--}2\text{ d}^{-1}$, significantly faster than bacterioplankton (Laws et al., 1987), as expected if one considers that in such systems the heterotrophic biomass exceeds that of autotrophs. In oligotrophic systems phytoplankters exhibit a dormant behaviour where photosynthesis rates are minimum, relying on a chance encounter with nutrient patches to revive their photosynthetic mechanism (Smetacek and Pollehne, 1986). In addition the maximum N:C and P:C ratios for phytoplankton have been altered to restrict the luxury uptake of nutrients.

In the microzooplankton module some food preference factors have been changed to account for the fact that dinoflagellates, unlike standard ERSEM V.11 where they were considered inedible, are parameterised as a food source for omnivorous microzooplankton. The bacterial specific uptake rate was significantly lowered, in order to reduce the bacterial growth rate in the range of $0.006\text{--}0.086\text{ (d}^{-1}\text{)}$, as suggested by Turley et al. (2000) and Pedros-Alio et al. (1999). Finally the assimilation efficiency was also lowered, simulating an average bacterial growth efficiency of 5–20% (Carlson and

Table 1. Parameters of the phytoplankton functional groups. P1 = diatoms, P2 = nanoalgae, P3 = picoalgae, and P4 = dinoflagellates. The parameters that have been changes from the standard ERSEM version 11 are indicated in bold. Parameter names used follow the nomenclature described in Blackford and Radford (1995)

Parameter	Name	P1	P2	P3	P4
Environmental effects					
Characteristic Q10	q10ST\$	2.0	2.0	2.0	2.0
Uptake					
max. specific upt at 10° C	sumST\$	1.0	1.2	1.3	1.0
Loss rates					
Excreted fraction of uptake	pu_eaST\$	0.05	0.2	0.2	0.05
Nutrient-lysis rate	sdoST\$	0.05	0.05	0.05	0.05
Nutrient-lysis rate under Si limitation	sdoST\$	0.1	–	–	–
Respiration					
Rest respiration at 10° C	srsST\$	0.15	0.1	0.1	0.1
Activity respiration	pu_raST\$		0.25	0.25	0.25
Nutrient Dynamics					
Min N/C ratio (mol gC ⁻¹)	qn1ST\$	0.00687	0.00687	0.00687	0.00687
Min P/C ratio	qn1ST\$	0.4288E-3	0.4288E-3	0.4288E-3	0.4288E-3
Redfield N/C ratio	qnRST\$	0.0126	0.0126	0.0126	0.0126
Redfield P/C ratio	qnRST\$	0.7862E-3	0.7862E-3	0.7862E-3	0.7862E-3
Multiplic fact min N/C ratio	xqcSTn\$	1.0	1.0	1.0	1.0
Multiplic fact min P/C ratio	xqcSTp\$	1.0	1.0	1.0	1.0
Multiplic fact max. N/C ratio	xqnST\$	2	2	2	2
Multiplic fact max. P/C ratio	xqpST\$	2	2	2	2
Max. Si/C ratio	qsSTc\$	0.03	–	–	–
Affinity for NO ₃	quSTn3\$	0.0025	0.0025	0.0025	0.0025
Affinity for NH ₄	quSTn4\$	0.0025	0.0025	0.0025	0.0025
Affinity for P	quSTp\$	0.0025	0.0025	0.0025	0.0025
Half value of Si limitation	chSTs\$	0.3	–	–	–
Sedimentation					
Nutrient limitation value for sedimentation	esNIST\$	0.7	0.75	0.75	0.75
Sinking rate (m day ⁻¹)	resSTm\$	5.0	0.0	0.0	0.0

Ducklow, 1996; Kirchman et al., 1991; Pedros-Alio et al., 1999; Turley et al., 2000).

3 Results and discussion

3.1 Hindcast experiments

3.1.1 Temperature

Temperature simulations plotted against in situ measurements for the 40, 60, 90 and 150 m depths are shown in Fig. 3. Stratification started approximately at the middle of April, reaching the highest subsurface values towards the end of October, and the location of the thermocline lay between 40 and 65 m. From November onwards a gradual mixing of the thermocline into deeper layers occurred until it disappeared in early January (Nittis et al., 2003). The bottom of the seasonal thermocline was at 50–70 m and, thus, the signal of the annual cycle of heating and cooling is hardly detected below that depth. At 40 m the model results follow very closely the field values, with minimum temperatures in March and Maximum in October. The large variability in

the in situ values of almost 4°C within a few days after June 2000 may be associated with changes in the eddy structure of the region. While the variation during October 2000 and December 2000 to January 2001 was coincident, with significant movements of the mooring line (Nittis et al., 2003). At 65 m the simulated temperature follows the general trend in the data, but fails to reproduce the observed variability. This variability is less pronounced than at 40 m and can be ascribed to similar processes. At both 90 m and 150 m there is no significant seasonality in the data and the model reproduces the observations.

3.1.2 In situ validation

The in situ data collected during the maintenance trips was compared with model results for nutrients and chlorophyll. Figure 4 shows simulated annual mean values plotted against depth, along with mean in situ concentrations for the period from 30 January 2000 until 19 April 2001, while range bars indicate minimum and maximum measured values. Model results are generally in good agreement with the observed in situ data in the whole water column, although the field data exhibits a substantial variability in almost all parameters.

Table 2. Parameters of microzooplankton functional groups and bacteria. Z5 = microzooplankton, Z6 = heterotrophic flagellates and B1 = bacteria. The parameters that have been changed from the standard ERSEM version 11 are indicated in bold. Parameter names follow the nomenclature described by Blackford and Radford (1995)

Parameter	Name	B1	Z6	Z5
Environmental effects				
Characteristic Q10	q10ST\$	2.95	2.0	2.0
Half oxygen saturation	chrSTo\$	0.3125	7.8125	7.8125
Uptake				
Half saturation value	chuSTc\$	30.0	250	200
Max. spec uptake rate 10° C	sumST\$	0.8	5.0	1.2
Availability of P1 for ST	suP1_ST\$	–	0.0	1.0
Availability of P2 for ST	suP2_ST\$	–	0.0	0.4
Availability of P3 for ST	suP3_ST\$	–	0.4	0.0
Availability of P4 for ST	suP4_ST\$	–	0.0	1.0
Availability of Z5 for ST	suZ5_ST\$	–	0.0	1.0
Availability of Z6 for ST	suZ6_ST\$	–	0.2	1.0
Availability of B1 for ST	suB1_ST\$	–	1.0	0.0
Selectivity	mindfoodST\$	–	100	30
Loss rates				
Assimilation efficiency	puST\$	0.25	0.4	0.5
Assimilation efficiency at low temp	puST0\$	0.2	–	–
Excreted fraction of uptake	pu_eaST\$	–	0.5	0.5
Excretion				
Fraction of excretion production to DOM	pe_R1ST\$ –	0.5	0.5	
Mortality				
Oxygen dependent mortality rate	sdST0\$	–	0.25	0.25
Temperature independent mortality	sdST\$	0.001	0.05	0.05
Respiration				
Rest respiration at 10° C	srsST\$	0.01	0.02	0.02
Nutrient Dynamics				
Max. N/C ratio	qnSTc\$	0.0208	0.0167	0.0167
Max. P/C ratio	qpSTc\$	0.00208	0.00167	0.00167

Field data (phosphate, nitrate, ammonia and silicate) indicate the presence of a weak nutracline at 80 m, which is successfully reproduced, although more pronounced in the model results. In the top 100 m the model phosphate values are close to the minimum measured concentrations, with the data fit improving below 100m. It should be noted that compared to the CINCS data set for the same region (Tselepides et al., 2000) (April 1994 – July 1996), phosphate values during the M3A deployment period are significantly higher. Nitrate exhibits a similar picture to phosphate with model concentrations close to the lower end of the range of in situ data. Once again, M3A nitrate concentrations are higher than the CINCS data and exhibit significantly higher variability at all depths. Simulated silicate and ammonia concentrations lie within the variability of measured concentrations. The maximum measured chl-a values at 80m are indicative of the deep chlorophyll maximum (DCM), a characteristic of the area (Tselepides et al., 2000), which is reproduced by the model at a slightly shallower depth. This is due to the fact that in the model the DCM coexists with the maximum biomass, while the analysis of the in situ data shows that the DCM is deeper compared to the biomass or cell count maximum

(Psarra et al., 2000), indicating an elevated chlorophyll per cell content at deeper layers. The model fails to reproduce chlorophyll in the deeper parts of the water column below 10 m, where small chlorophyll levels are detected even down to 200–400 m due to wind mixing, as has been suggested by Tselepides and Polychronaki (1996).

3.1.3 Chlorophyll

Figure 5 shows the simulated chl-a against the in situ fluorometer derived concentrations. Model results are very close to the measured concentrations at 40 m, with the exception of the period from 22 August 2000 until 29 October 2000, which can be attributed to biofouling of the sensor and the vertical displacement of the mooring line (Nittis et al., 2003). During this period the comparison of model and data is meaningless. This argument is strengthened by the fact that after 31 October 2000, the measured chl-a concentrations are once more very close to the model simulation.

At 65 m the model overestimated chlorophyll during March and April 2000, producing a peak at the end of March due to intense mixing. For the rest of the deployment period the simulation is very good, closely following the measured

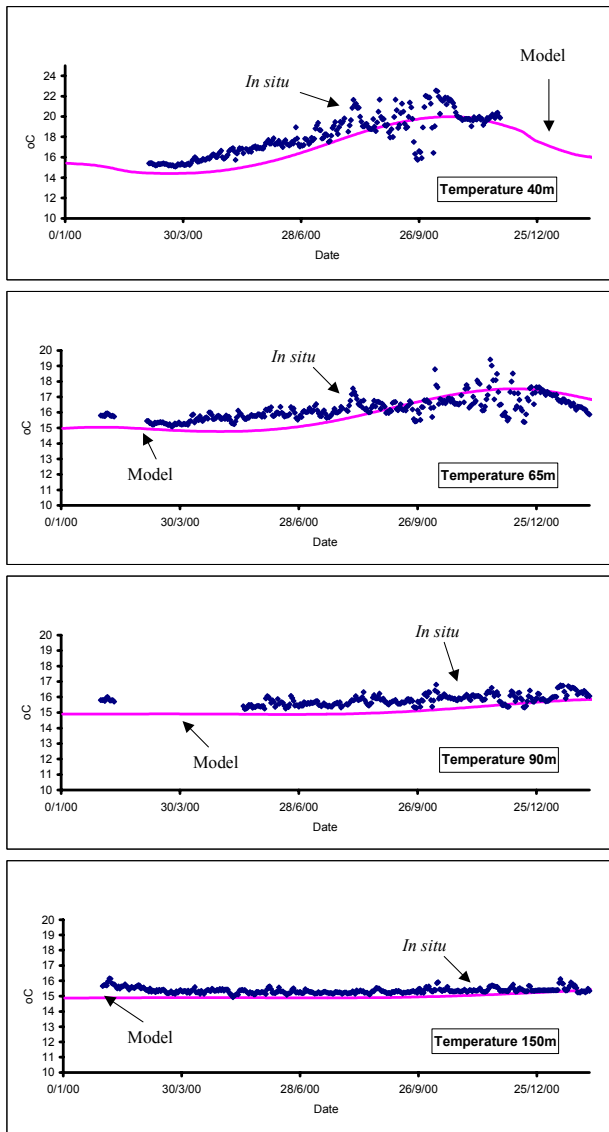


Fig. 3. Validation of model temperature with CTD measurements at 40, 65, 90 and 150 m.

concentrations, producing a peak in October and minimum values in February and August 2000.

At 90 m field data was unavailable for the first seven months due to sensor problems. For the rest of the deployment period the model, although following the measured trend, systematically produced lower concentrations. Analogous model behaviour is observed at 115 m, where there is an underestimation of Chl-a, although differences appear significant only during the first seven months. This poor behaviour of the model may partly be due to the use of a constant C:Chl ratio at all depths. It is well documented that there is a common trend of an increase in chlorophyll-a and other light harvesting pigments as growth irradiance decreases (Dubinsky, 1992; Estrada, 1985; Estrada et al., 1993; Gasol et al., 1997; Venrick, 1990). However, there is a rather wide range of C:Chl values suggested by researchers for different environ-

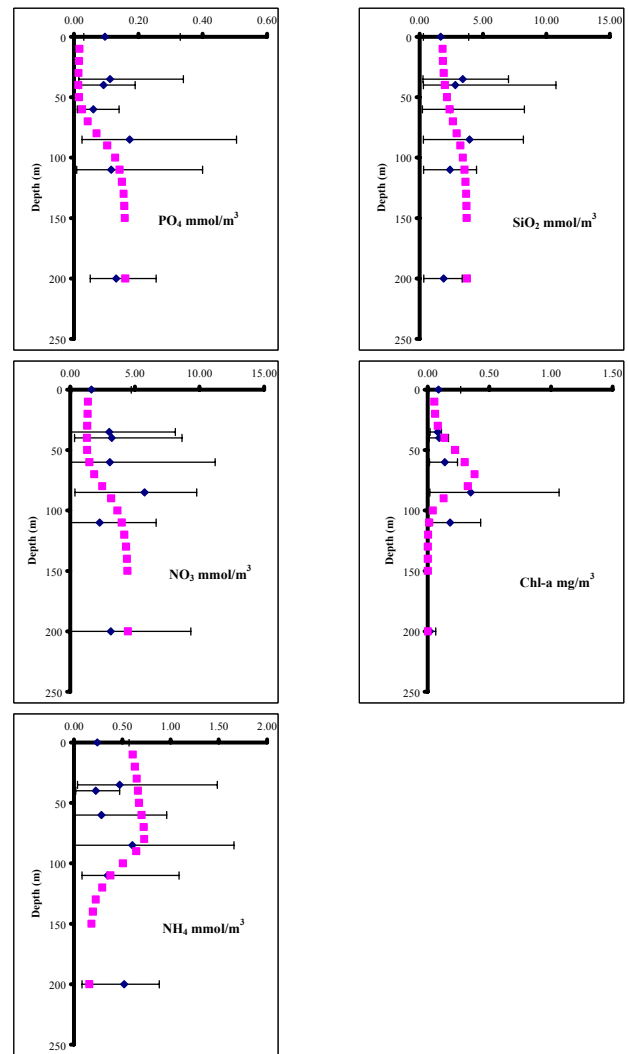


Fig. 4. Validation of model Phosphate, Nitrate, Ammonia, Silicate and Chl-a with in situ data obtained during maintenance trips. All sampling measurements at a particular depth are treated as replicates and averaged. Range bars are used to indicate variability (minimum and maximum values).

ments, which confuses the parameterisation issue. A variable C/Chl ratio of decreasing values with increasing depth would improve the simulations at the deeper layers. An additional further source of discrepancy may arise from the calibration of the data, since there is a wide variation even within three days in the in situ measurements.

3.2 Oxygen

The dissolved oxygen sensors only gave reliable data during the first six months of the buoy deployment and then failed at the beginning of July 2000 (Fig. 6). At 40 m model results generally underestimate the oxygen concentration, except at the end of March 2000. Since the region is one of low productivity, this discrepancy can probably be ascribed to the fact that the model overestimates temperature at that depth. A

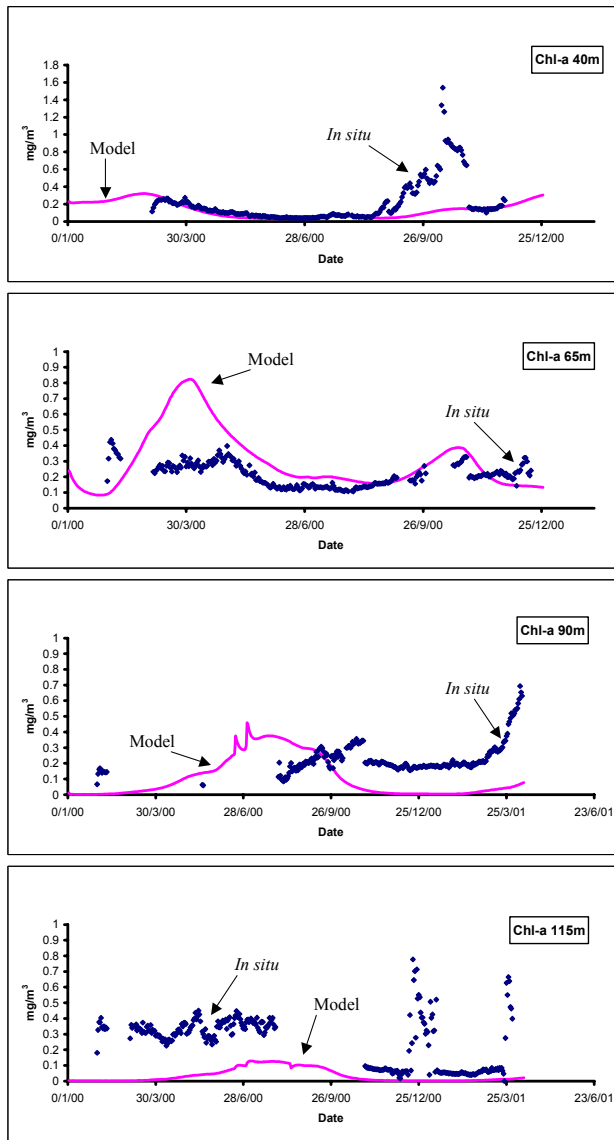


Fig. 5. Validation of model chlorophyll concentration with CTD measurements at 40, 65, 90 and 115 m.

better agreement between simulation and field data is found at 65 m and 115 m, where the temperature simulations have a better fit with data. At 65 m the maximum oxygen concentrations are found during April–May, which may be attributed to increased primary production, as evidenced by the Chl-a concentrations.

3.2.1 Nitrate

Although model results (Fig. 7) are within the right range of measured values, they do not closely follow the in situ variability. The extreme high values measured at the beginning of the data set are due to an unsuccessful deployment during which the analyser was almost at the surface. The marked decrease in nitrate during mid-May, suggesting an increased activity of primary producers, cannot be explained by the chl-

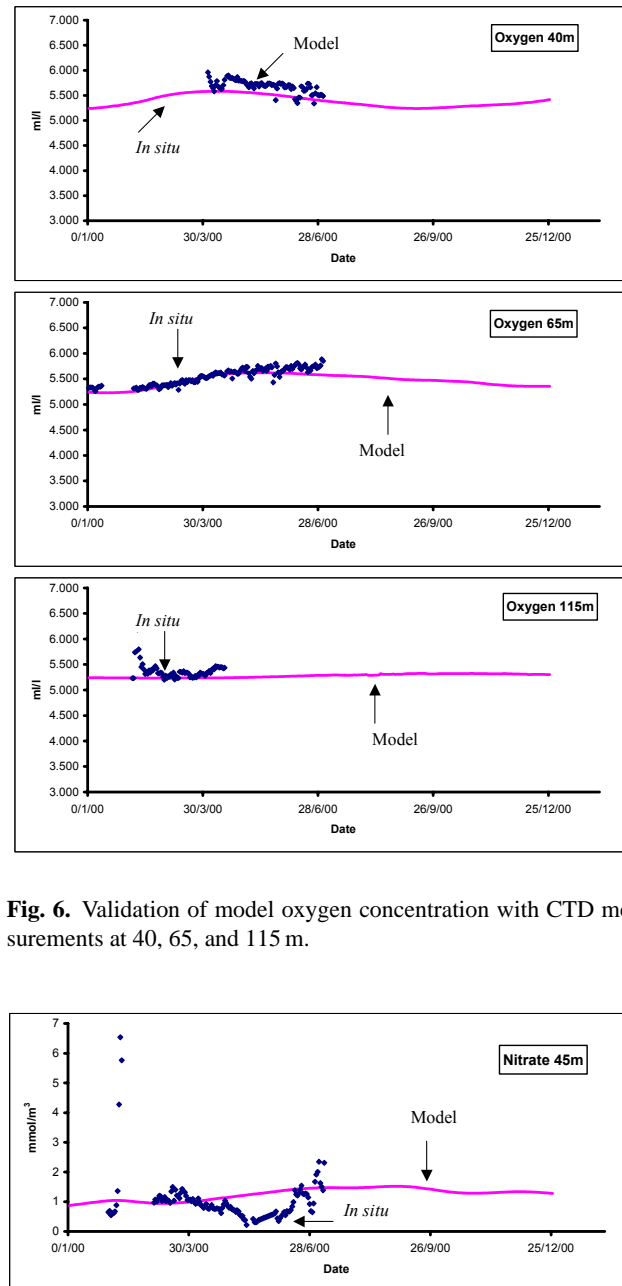


Fig. 6. Validation of model oxygen concentration with CTD measurements at 40, 65, and 115 m.

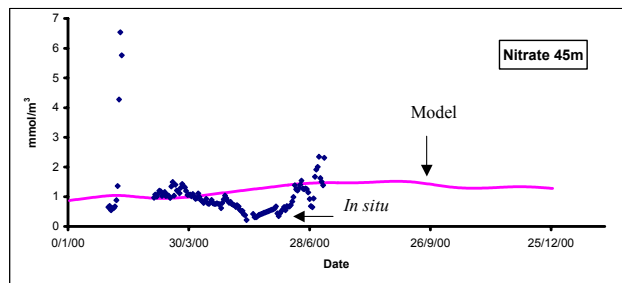


Fig. 7. Validation of model nitrate concentration at 45 m, comparison of nitrate concentrations from nitrate analyzer and in situ measurements during maintenance trips.

a measurements at the 40 and 65 m layers, where concentrations are either unchanged or decreasing. The failure of the model to reproduce this effect can be attributed to the lack of lateral advection of nitrate through the system. The observed nitrate concentrations in winter are lower than might be expected from previous studies. It should be noted that strong mixing events during winter were not observed during the deployment period. It has been suggested by Georgopoulos et al. (1989); Theocharis (1995); Theocharis et al. (1993) that such events may significantly affecting the biology of the

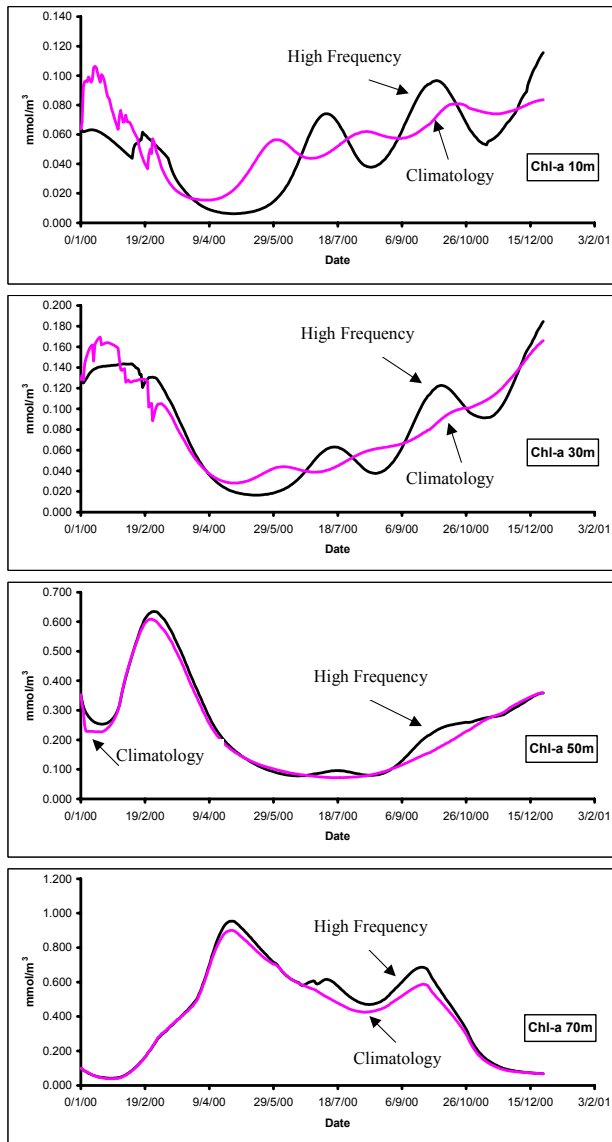


Fig. 8. Model chlorophyll simulations at various depths with climatological and real-time forcing.

area.

3.3 Sensitivity analysis, the response of the model to changes in forcing

The effects of changes in the atmospheric forcing upon the model behaviour were investigated by making two separate runs. In the first run, climatological wind forcing was used and the model was relaxed to climatological sea surface temperature, and salinity obtained from the Mediterranean Ocean Data Base database (Brasseur et al., 1996). For the second run, high-frequency wind forcing (half hour), for the period from 1 January 2000 until 3 February 2001, for the specific area, was obtained from the POSEIDON system, while the model was relaxed to mean monthly in situ sea surface temperature and salinity. The simulations of chloro-

phyll at the depths of 10, 30, 50 and 70 m from the two runs are shown in Fig. 8.

In the deeper layers, at 50 and 70 m, there are no significant differences between the two runs, and the model produces a spring and autumn bloom. At 50 m the spring bloom is observed during March, while at 70 m the spring bloom is observed during the end of April/beginning of May. In both layers the autumn bloom is shown during September–October.

However, in the upper layers (10 and 30 m), there are significant differences between the two runs, with more pronounced minimum and maximum chlorophyll concentrations in the run with the high-frequency data forcing, due to the diffusion of nutrients caused by the vertical mixing processes, which are much better described.

Thus, the model results indicate that the 1-D physical model only responds to sea surface forcing in the near surface layers and that the effect of high-frequency forcing data in the simulation of the ecosystem is only significant in the mixed layer. As mentioned above, the appearance of the Transition Mediterranean Water (TMW) at approximately 300–500 m, characterized by high nutrient and low oxygen concentrations in conjunction with the gyral features populating the area, significantly affects the intermediate layers of the Cretan Sea. Thus, in the deeper layers of the euphotic zone, the ecosystem functioning is significantly affected by the advective processes, which cannot be described by the present model requiring a 3-D model.

3.4 Model performance

The performance of the model is evaluated using the misfit error between modelled temperature (Fig. 9), Chl-a concentrations (Fig. 10), and in situ data. The model underestimates the temperature at 40 m by around 10–15% for most of the year, except in the autumn, when it is overestimated. At 65 m a similar trend is observed but the errors are smaller (5–10%). The model is consistently colder than the observations, which implies that the buoyancy is weaker than reality and that vertical mixing may be overestimated. The implementation of a surface heat flux formulation in the model may help to overcome these problems, as it would place the ecosystem model in a fully 3-D hydrodynamic environment. The data mismatch of chlorophyll is shown in Fig. 10. At 40 m the model consistently underestimates the data by 50–100%, except during July, while at 65 m Chl-a is consistently overestimated by 50–100%, except during the bloom period, when the error is larger. There are a number of possible reasons for this discrepancy. The attenuation of light down the water column may be incorrect, and as previously, noted the vertical mixing may be wrong. Alternatively, the use of a fixed carbon to chlorophyll ratio in the model may cause this difference, with the ratio being underestimated in the near surface water and overestimated towards the bottom of the euphotic zone. The absence of phytoplankton biomass and primary production measurements precludes us from making further conclusions. However, it seems apparent that further

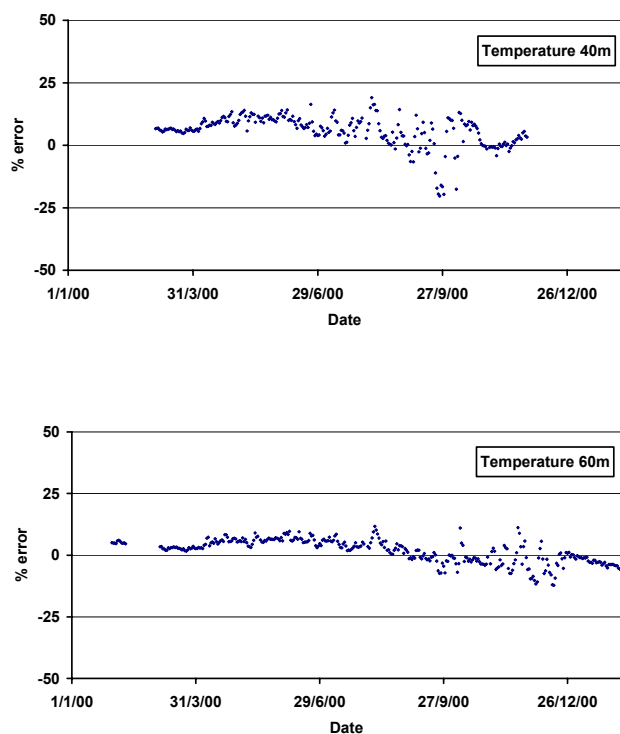


Fig. 9. The percentage error ($100 \cdot (\text{Data} - \text{Model}) / \text{Data}$) for temperature at (a) 40 m and (b) 60 m.

work is required to improve the response of the phytoplankton model to low light environments. One possible solution is to implement a variable carbon-chlorophyll model, such as that of Geider et al. (1997).

The fit with nitrate at 45 m is shown in Fig. 11. In situ data and the model show a good fit in March and June/July, but overestimates the concentrations by 200–300% in between. The excess of nitrate in the spring may arise from the lack of horizontal advective processes in the model or through problems with model phytoplankton uptake of nutrients and nutrient recycling mechanisms. Oxygen data (not shown) underestimates the data by up to 5% at 40 m, 65 m and 115 m. This reinforces the previous conclusion that in a low productivity environment, the oxygen concentration is primarily temperature dependent.

Alternatively, data fits can be substantially improved by the implementation of a data assimilation scheme, such as the Ensemble Kalman Filter. This has been demonstrated for the M3A data set by Allen et al. (2003).

4 Conclusions

For the first time the oligotrophic ecosystem of the Cretan Sea has been monitored over an extended period of time, providing valuable information and a unique opportunity for ecosystem model application. The Cretan Sea ecosystem model developed during the MATER project when forced with real atmospheric data simulates very satisfactorily the

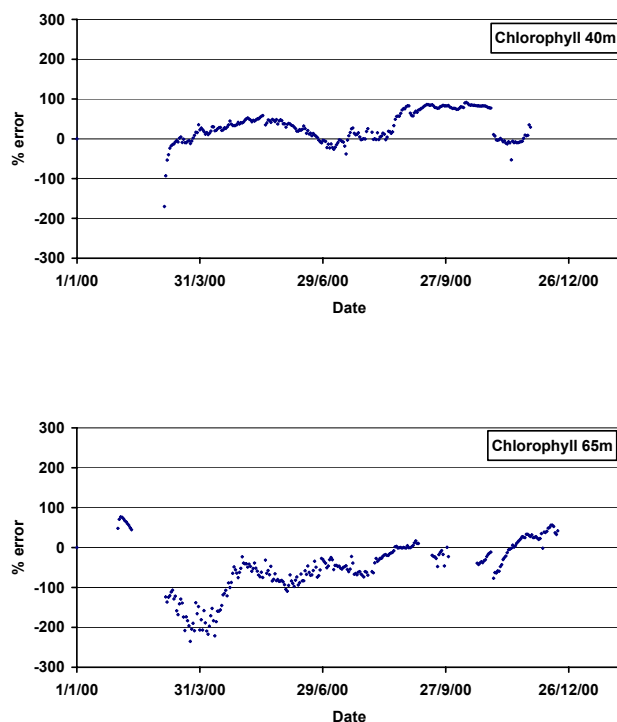


Fig. 10. The percentage error ($100 \cdot (\text{Data} - \text{Model}) / \text{Data}$) for chlorophyll-a at (a) 40 m and (b) 60 m.

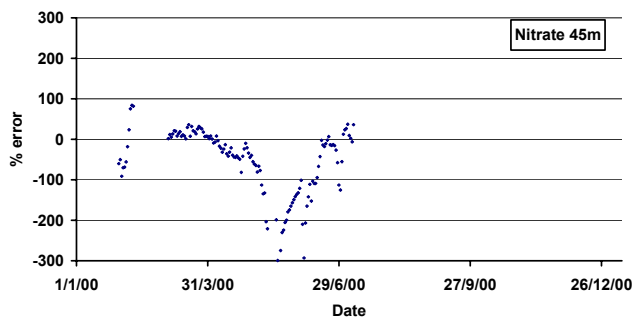


Fig. 11. The percentage error ($100 \cdot (\text{Data} - \text{Model}) / \text{Data}$) for nitrate at 45 m.

chl-a time series in the upper part of the water column as recorded by the fluorometers. Deeper down, close to the DCM the underestimation of the model concentrations can be attributed to the elevated cell chlorophyll content, a physiological adaptation to lower light conditions. A future modification of the ecosystem model to account for this phenomenon, as well as the implementation of heat fluxes computed for the particular area, are considered necessary.

Although the in situ nitrate data are rather incomplete as a result of instrument malfunction such data, are, however, very important in these studies. Considering the phosphate limitation of the area high-frequency phosphate data from an autonomous instrument would provide significant information towards the improvement of the model.

Laboratory calibration of the fluorometric sensors is considered crucial and necessary, in addition to the calibration with in situ data due to the significant variability of the chlorophyll concentrations in the samples in short-time periods (2–3 days).

Due to its 1-D nature, the model does not include important processes of the particular area, such as horizontal advection and vertical mixing arising from circulation patterns. Nevertheless, it provides a very successful numerical base for further development towards a 3-D forecasting ecosystem model and the implementation of assimilation techniques.

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References

- Abbott, M. R.: Phytoplankton patchiness: ecological implications and observation methods. In: *Patch Dynamics, Lecture Notes in Biomathematics*, (Eds) Levin, S. A., Powell, T. M., and Steele, J. H., Springer Verlag, Berlin Heidelberg, 307, 1993.
- Allen, J. I., Blackford, J. C., and Radford, P. J.: An 1-D vertically resolved modelling study of the ecosystem dynamics of the middle and southern Adriatic Sea, *J. Marine Systems*, 18, 265–286, 1998.
- Allen, J. I., Somerfield, P. J. and Siddorn, J. R.: Primary and bacterial production in the Mediterranean Sea: a modelling study, *J. Marine Systems*, (in press), 2002.
- Allen, J. I., Ekenes, M. and Evensen, G.: An Ensemble Kalman Filter with a complex marine ecosystem model: Hindcasting phytoplankton in the Cretan Sea, *Ann. Geophysicae*, this issue, 2003.
- Azov, Y.: Eastern Mediterranean – a Marine Desert? *EMEC'S'90*, 23, 225–232, 1991.
- Balopoulos, E. T.: PELAGOS, MAS2-CT93-0059, NCMR, Athens, 1996.
- Baretta, J. W., Ebenhoh, W., and Ruurdij, P.: The European Regional Seas Ecosystem Model, a complex marine ecosystem model, Netherlands, *J. Sea Research*, 33, 233–246, 1995.
- Baretta-Bekker, J. G., Baretta, J. W., Hansen, A. S., and Riemann, B.: An improved model of carbon and nutrient dynamics in the microbial food web in marine enclosures, *Aquatic microbial ecology*, 14, 91–108, 1998.
- Baretta-Bekker, J. G., Baretta, J. W., and Rasmussen, E.: The microbial foodweb in the European regional Seas Ecosystem Model, Netherlands, *J. Sea Research*, 33, 363–379, 1995.
- Becacos-Kontos, T.: Primary production and environmental factors in an oligotrophic biome in the Aegean Sea, *Marine Biology*, 42, 93–98, 1977.
- Blackford, J. and Radford, P.: A structure and methodology for Marine Ecosystem Modelling, Netherlands *J. Sea Res.*, 33, 247–260, 1995.
- Berland, B., Bonin, D., and Maestrini, S.: Azote ou phosphore? Considerations sur le “paradoxe nutritionnel” de la Mer Méditerranée, *Oceanologica Acta*, 3, 135–142, 1980.
- Blumberg, A. F. and Mellor, G. L.: A Coastal Ocean Numerical Model. In: *Mathematical Modelling of Estuarine Physics*, (Eds) Sunderman, J. and Holtz, K., Proceedings of the International Symposium, Springer-Verlag Berlin, Hamburg, 203–214, 1978.
- Brasseur, P., Brankart, J.-M., Schoenauen, R., and Beckers, J.-M.: Seasonal Temperature and Salinity Fields in the Mediterranean Sea: Climatological Analyses of an Historical Data Set, *Deep Sea Res.*, 43, 159–192, 1996.
- Carlson, C. A. and Ducklow, H. W.: Growth of bacterioplankton and consumption of dissolved organic carbon in the Sargasso Sea, *Aquatic Microbial Ecology*, 10, 69–85, 1996.
- Dubinsky, Z.: The functional and optical absorption cross-sections of phytoplankton photosynthesis. In: *Primary Productivity and Biogeochemical Cycles in the Sea*, (Eds) Falkowski, P. G. and Woodhead, A. D., Plenum Press, New York, 550, 1992.
- Estrada, M.: Primary production at the deep chlorophyll maximum in the western Mediterranean. In: *19th European Marine Biology Symposium*, (Ed) Gibbs, P. E., 109–121, 1985.
- Estrada, M., Marrase, C., Latasa, M., Berdalet, E., Delgado, M., and Riera, T.: Variability of deep chlorophyll maximum characteristics in the Northwestern Mediterranean, *Marine Ecology Progress Series*, 92, 289–300, 1993.
- Gasol, J. M., del Giorgio, P. A., and Duarte, C. M.: Biomass distribution in marine planktonic communities, *Limnology Oceanography*, 42, 6, 1353–1363, 1997.
- Geider, R. J., MacIntyre, H. L., and Kana, T. M.: A dynamic growth model of phytoplankton growth and acclimation: responses of the balanced growth rate and chlorophyll-a: carbon ratio to light limitation and temperature, *Mar. Ecol. Prog. Ser.*, 148, 187–200, 1997.
- Georgopoulos, D., Theocharis, D., and Zodiatis, G.: Intermediate water formation in the Cretan Sea (S. Aegean Sea), *Oceanologica Acta*, 12, 4, 353–359, 1989.
- Gotsis-Skretas, O., Pagou, K., Moraitou-Apostolopoulou, M., and Ignatiades, L.: Seasonal horizontal and vertical variability in primary production and standing stocks of phytoplankton and zooplankton in the Cretan Sea and the straits of the Cretan Arc (March 1994–January 1995), *Progress in Oceanography*, 44, 625–649, 1999.
- Hellermann, S. and Rosenstein, M.: Normal wind stress over the world ocean with error estimates. *J. Phys. Oceanography*, 13, 1093–1104, 1983.
- Ignatiades, L., Georgopoulos, D., and Karydis, M.: Description of the phytoplanktonic community of the oligotrophic waters of the SE Aegean Sea (Mediterranean), *P.S.Z.N. I: Marine Ecology*, 16, 1, 13–26, 1995.
- Kirchman, D. L., Suzuki, Y., Garside, C., and Ducklow, H. W.: High turnover rates of dissolved organic carbon during a spring phytoplankton bloom, *Nature*, 352, 612–614, 1991.
- Krom, M. D., Brenner, S., Kress, N., Neori, A., and Gordon, L. I.: Nutrient dynamics and new production in a warm-core eddy from the Eastern Mediterranean Sea, *Deep-Sea Res.*, 39, 3/4, 467–480, 1992.
- Krom, M. D., Brenner, S., Kress, N., Neori, A., and Gordon, L. I.: Nutrient distributions during an annual cycle across a warm-core eddy from the E. Mediterranean Sea, *Deep-Sea Res.*, 40, 4, 805–825, 1993.
- Krom, M. D., Kress, N., and Brenner, S.: Phosphorus limitation of primary productivity in the eastern Mediterranean Sea, *Limnology Oceanography*, 36, 3, 424–432, 1991.
- Laws, E. A., DiTullio, G. R. and Redalje, D. G.: High phytoplankton growth and production rates in the North Pacific subtropical

- gyre, *Limnology Oceanography*, 32, 905–918, 1987.
- Mellor, G. L. and Yamada, T.: Development of a Turbulence Closure Model for Geophysical Fluid Problems, *Review Geophysics and Space Physics*, 20, 851–875, 1982.
- Nittis, K., Tziavos, C., Thanos, I., et al.: The Mediterranean Moored Multi-sensor Array (M3A): System Development and Initial Results, *Ann. Geophysicae*, this issue, 2003.
- Patsch, J.: MACADOB a model generating synthetic time series of solar radiation for the North Sea, *Ber. Zentrum Meeresforsch. Klimaforsch.*, B 16, 1–67, 1995.
- Pedros-Alio, C., Calderon-Paz, J. I., Guixa-Boixereu, N., Estrada, M., and Gasol, J. M.: Bacterioplankton and phytoplankton biomass and production during summer stratification in the northwestern Mediterranean Sea, *Deep-Sea Res. I*, 46, 985–1019, 1999.
- Psarra, S., Tselepides, A., and Ignatiades, L.: Primary productivity in the oligotrophic Cretan Sea (NE Mediterranean): seasonal and interannual variability, *Progress in Oceanography*, 46, 187–204, 2000.
- Smetacek, V. and Pollehne, F.: Nutrient cycling in pelagic systems: a reappraisal of the conceptual framework, *Ophelia*, 26, 401–428, 1986.
- Soukissian, T., Chronis, G., and Nittis, K.: POSEIDON: operational marine monitoring system for Greek Seas, *Sea Technology*, 40, 31–37, 1999.
- Stergiou, K. I., Christou, E. D., Georgopoulos, D., Zenetos, A., and Souvermezoglou, C.: The Hellenic Seas: Physics, Chemistry, Biology and Fisheries, *Oceanography and Marine Biology*, 35, 415–538, 1997.
- Theocharis, A.: Hydrological and dynamical structure of the South Aegean Sea: Flows through the straits of the Cretan Arc, Athens, 1995.
- Theocharis, A., Balopoulos, E., Kioroglou, S., Kontoyiannis, H., and Iona, A.: A synthesis of the circulation and hydrography of the South Aegean Sea and the Straits of the Cretan Arc (March 1994–January 1995), *Progress in Oceanography*, 44, 469–509, 1999.
- Theocharis, A., Georgopoulos, D., Lascaratos, A., and Nittis, K.: Water masses and circulation in the central region of the Eastern Mediterranean: Eastern Ionian, South Aegean and Northwest Levantine, *Deep-Sea Res. II*, 40, 6, 1121–1142, 1993.
- Thingstad, T. F. and Rassoulzadegan, F.: Nutrient limitations, microbial food webs, and “biological C-pumps”: suggested interactions in a P-limited Mediterranean, *Marine Ecology Progress Series*, 117, 299–306, 1995.
- Triantafyllou, G., Petihakis, G., Allen, J. I., and Tselepides, A.: Primary Production and Nutrient Dynamics of the Cretan Sea Ecosystem (North Eastern Mediterranean), A Modelling Approach, *J. Marine Research*, submitted, 2002.
- Tselepides, A. and Polychronaki, T.: Pelagic-benthic coupling in the oligotrophic Cretan Sea (NE Mediterranean), *IMBC*, Iraklio, 1996.
- Tselepides, A., Zervakis, V., Polychronaki, T., Donavaro, R., and Chronis, G.: Distribution of nutrients and particulate organic matter in relation to the prevailing hydrographic features of the Cretan Sea (NE Mediterranean), *Progress in Oceanography*, 46, 113–142, 2000.
- Turley, C. M., Bianchi, M., Christaki, U., Conan, P. Harris, J. R. W., Psarra S., Ruddy, G., Stutt, E. D., Tselepides, A., and van Wambeke, F.: Relationship between primary producers and bacteria in an oligotrophic sea – the Mediterranean and biogeochemical implications, *Marine Ecology Progress Series*, 193, 11–18, 2000.
- Venrick, E. L.: Phytoplankton in an oligotrophic ocean: species structure and interannual variability, *Ecology*, 71, 4, 1547–1563, 1990.
- Zavatarelli, M., Barreta, J. W., Barreta-Bekker, J. G., and Pinardi, N.: The dynamics of the Adriatic Sea ecosystem, An idealized model study, *Deep Sea Res. I*, 47, 5, 937–970, 2000.